

## **Manipulating photorespiration to increase plant productivity: recent advances and perspectives for crop improvement.**

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1    **ABSTRACT**

2    Recycling of the 2-phosphoglycolate generated by the oxygenase reaction of  
3    Rubisco requires a complex and energy-consuming set of reactions collectively  
4    known as the photorespiratory cycle. Several approaches have been proposed  
5    with the aim of producing plants with reduced rates of photorespiration energy or  
6    carbon loss, both by screening for natural variation and by means of genetic  
7    engineering. Recent works indicate that plant yield can be substantially improved  
8    by the alteration of photorespiratory fluxes or by engineering artificial bypasses  
9    to photorespiration. However, there is also evidence indicating that, under certain  
10   environmental and/or nutritional conditions, reduced photorespiratory capacity  
11   may be detrimental for plant performance. Here, we summarize recent advances  
12   obtained in photorespiratory engineering and discuss prospects for these advances  
13   to be transferred to major crops to help address the globally increasing demand  
14   for food and biomass production.

15  
16   **Keywords**

17   Crops, Food production, Genetic engineering, Photorespiration, Rubisco, Yield  
18   improvement

19  
20   **Highlight**

21   Manipulation of the photorespiratory pathway may greatly increase plant  
22   productivity. Here we summarize recent advances in the engineering of  
23   photorespiration and discuss how to use these approaches for crop improvement.

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## Introduction

There is an urgent demand for increased crop productivity due to the world's population growth, increasing global affluence, reduction of cultivable soils and higher demand for plant based biofuels. The required increase in agricultural productivity required by 2030 may be in the range of 60 to 120% as compared to the levels of 2005 (Ort *et al.*, 2015). A rapid increase in crop yield, especially for cereals, was obtained in the second half of the 20<sup>th</sup> century during the so-called "Green Revolution". Resulting from breeding strategies, this led to the introduction of new crop strains with a greater proportion of biomass partitioned into grains and greater inputs of fertilizer, pesticides and water. However, increases in yield for several major crops such as rice in recent years have been scarce (Zhu *et al.*, 2010), and it is possible that actual crop yield is approaching the ceiling of maximal yield potential (Tilman *et al.*, 2002). Further increases in nitrogen and phosphorous fertilization are unlikely to solve this problem and indeed many countries are currently attempting to reduce the levels of fertilization used in intensive agriculture. For these reasons, attention is being paid to the improvement of photosynthesis, a process that is still far from its theoretical maximum efficiency. Several recent reviews summarise the opportunities that have been so far identified to improve photosynthetic efficiency (Zhu *et al.*, 2010; Raines, 2011; Maurino and Weber, 2013; Long *et al.*, 2015; Ort *et al.*, 2015).

Photosynthetic CO<sub>2</sub> fixation starts with the carboxylation of ribulose 1,5-bisphosphate (RuBP), catalysed by ribulose 1,5-bisphosphate carboxylase-oxygenase (Rubisco), to yield two molecules of 3-phosphoglycerate (3PGA). An unavoidable side reaction of Rubisco is the oxygenation of RuBP to produce one molecule of 3PGA and one molecule of 2-phosphoglycolate (2PG). Photosynthetic organisms evolved a complex pathway to recycle 2PG that involve reaction taking place in chloroplasts, peroxisomes, mitochondria and the cytosol, (Bauwe *et al.*, 2010). In this photorespiratory cycle, two molecules of 2PG are transformed into one molecule of 3PGA and one carbon atom is lost as CO<sub>2</sub> with an addendant cost of 4 NAD(P)H and 7 ATP. Photorespiration has long been viewed as a target for crop improvement due to the seemingly wasteful nature of the cycle and the high energetic cost that it imposes on plant metabolism.

The cost of photorespiration is massive at both the leaf and canopy scale. CO<sub>2</sub> is lost from photorespiration under 25°C at about 25% the rate of net CO<sub>2</sub> fixation (Sharkey, 1985; Sage *et al.*, 2012). For example, photorespiration results in the loss of ~322 trillion Calories annually in the US Corn Belt alone. Even a 5% reduction in photorespiration would be worth almost \$540 million a year in yield gain in this growing region (Walker *et al.*, submitted for publication). This high cost stems in part from the energy used in the reassimilation of the ammonia produced following glycine decarboxylation in the mitochondrion. Moreover, rates of photorespiration increase with temperature and the scarcity of water as these conditions favour increased Rubisco oxygenation (Walker *et al.*, submitted for publication). It is thus not surprising that several groups tried to develop plants with reduced rates of photorespiration with the aim of increasing productivity (Peterhänsel *et al.*, 2013a). However, the view of photorespiration as a pathway that only aims at recycling the carbon of 2PG may be simplistic. In addition to photosynthesis, photorespiration interacts with several central metabolic pathways (Foyer *et al.*, 2009; Bauwe *et al.*, 2010; Fernie *et al.*, 2013), and both the relevance and the regulatory aspects of these interactions need further investigations. Furthermore, photorespiration may contribute substantially to the production of serine (Benstein *et al.*, 2013; Ros *et al.*, 2013) and has been implicated in the response to certain biotic (Taler *et al.*, 2004) and abiotic stresses (Wingler *et al.*, 2000; Voss *et al.*, 2013). It was additionally recently demonstrated that there is a positive correlation between photorespiration and productivity (Aliyev, 2012) and between photorespiration and nitrate assimilation (Bloom *et al.*, 2010). While most efforts are aimed at generating plants with reduced photorespiratory rates, the eventual performance of these plants in the field and thus under stress conditions needs also to be considered. Tantalizing results have been obtained by re-engineering photorespiratory pathway in model plants (Kebeish *et al.*, 2007; Timm *et al.*, 2012a) or easy to transform non-staple crops such as tobacco (Lin *et al.*, 2014a), the transfer of these manipulations to our major crops and demonstration of benefits under field conditions is still lacking. In this article we summarise the different approaches that have been used to manipulate photorespiration and their possible application for crop improvement.

*Screening for plants with naturally reduced rates of photorespiration*

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103 Screenings of mutagenized plants that showed an altered phenotype under normal  
104 air conditions but not under conditions in which photorespiration is suppressed  
105 (CO<sub>2</sub>-enriched atmosphere) were carried in several C<sub>3</sub> species, notably barley and  
106 Arabidopsis (Sommerville and Ogren, 1992; Blackwell *et al.*, 1988; Foyer *et al.*,  
107 2009; Peterhänsel *et al.*, 2010). This approach permitted the identification of the  
108 genes that encode for the core enzymes of the photorespiratory cycle. However,  
109 the mutants obtained generally show poor performance under normal air  
110 conditions associated with different stress symptoms (Timm and Bauwe, 2013). In  
111 another approach, natural variants with reduced rates of photorespiration  
112 associated with higher yields were screened across broad populations. While  
113 preliminary trials carried out with tobacco gave promising results (Zelitch and  
114 Day, 1973), subsequent studies failed to identify plants with low levels of  
115 photorespiration paralleled by high productivity. Zelitch (1989) successfully  
116 isolated plants resistant to high levels of O<sub>2</sub> but the trait seemed more related to  
117 increased levels of catalase than to reduced rates of photorespiration. Other works  
118 of the same author identified tobacco plants with low photorespiratory rates and  
119 high catalase activity associated to higher yield, but this increase in yield was not  
120 robust across harvests (Brisson *et al.*, 1998; Zelitch, 1992). Similarly, screening of  
121 mutagenized tobacco plants identified genotypes with higher yield at low CO<sub>2</sub>  
122 concentrations but the high yield trait could not be related to reduced  
123 photorespiration (Medrano *et al.*, 1995). A more recent study that summarized the  
124 data obtained over 40 years of field trials using two major crop species, wheat and  
125 soybean, concluded that attempts to find highly productive genotypes with high  
126 photosynthetic but low photorespiratory rates are inconsistent instead showing  
127 that the highly productive cultivars have high rates of photosynthesis  
128 accompanied by high rates of photorespiration (Aliyev, 2012). These results,  
129 argue against the use natural variation as a strategy to alleviate the yield penalty of  
130 photorespiration suggesting that genetic engineering might be the only viable  
131 route.

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133 *Enhancing the amount of photorespiratory CO<sub>2</sub> scavenging*

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135 The CO<sub>2</sub> released during the decarboxylation step of photorespiration in  
136 mitochondria is not completely lost for the plant. On its way out of the cell, the  
137 released CO<sub>2</sub> can be refixed while passing through the chloroplasts (Sage and  
138 Sage R, 2009; Busch *et al.*, 2013). Some plants optimized this mechanism known  
139 as photorespiratory CO<sub>2</sub> scavenging by maximizing the likelihood for CO<sub>2</sub> to pass  
140 the chloroplasts. Firstly, these plants enhanced the surface of chloroplasts via  
141 stromules, connecting them to a net like structure (Sage and Sage R, 2009).  
142 Secondly, they associated chloroplasts tightly with mitochondria and peroxisomes  
143 (Sage and Sage R, 2009; Busch *et al.*, 2013). Rice has such morphological  
144 features and it was shown that its CO<sub>2</sub> compensation point is lower than that of  
145 other C<sub>3</sub> crops not showing this morphological adaption (Sage *et al.*, 2009).  
146 Similar to rice, the dicot C<sub>3</sub> plants *Flaveria pringlei* and *Flaveria robusta* also  
147 associated all three organelles and showed a reduced CO<sub>2</sub> compensation point  
148 compared to other C<sub>3</sub> *Flaveria* species (Sage *et al.*, 2013; Sage *et al.*, 2014).  
149 Although the effect of this anatomical adaption is not as big as the one found in  
150 C<sub>4</sub> or C<sub>2</sub> photosynthesis plants, it still accounts as a considerable improvement  
151 (Sage *et al.*, 2013). Therefore, installing this anatomy in a C<sub>3</sub> crop plant might be  
152 an alternative approach to optimize the yield. Compared to other approaches, a  
153 modification of cell anatomy should have little impact on cells metabolism. To  
154 install this anatomy in a plant, a better understanding of organelle movement and  
155 partitioning is needed. Natural varieties of rice and other plants showing an  
156 enhanced chloroplast surface and tight connecting of the three organelles should  
157 be analysed. Additionally a mutant screen of these varieties combined with RNA  
158 sequencing might reveal major regulators for the anatomy of cell organelles.  
159 Interestingly, in *Arabidopsis thaliana*, it was shown that stromules, which are  
160 used to enlarge the chloroplast surface, were established when plants were  
161 stressed with heat (Holzinger *et al.*, 2007). It would therefore be of interest to  
162 study mutant lines affected in stromule formation such as *arc(s)* (Holzinger *et al.*,  
163 2008), or even lines affected in chloroplast movement such as *chup1* (Oikawa *et*  
164 *al.*, 2008) and compare the rates of CO<sub>2</sub> fixation of these mutants with the wild-  
165 type ones.

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167 *Introducing C<sub>4</sub> metabolism into C<sub>3</sub> species*

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C<sub>4</sub> photosynthesis greatly reduces photorespiration by concentrating CO<sub>2</sub> at the active site of Rubisco. With the exception of the so-called single-cell C<sub>4</sub> plants (Sharpe and Offermann, 2014), C<sub>4</sub> plants have adopted different biochemical and anatomical modifications. C<sub>4</sub> leaves have two distinct layers of photosynthetic tissue (the so called “Kranz” leaf anatomy): mesophyll cells that are in contact with atmospheric CO<sub>2</sub> via intercellular air spaces, and bundle sheath cells with cell walls that are less permeable to CO<sub>2</sub>. CO<sub>2</sub> is assimilated into oxaloacetate in the mesophyll cells via PEP carboxylase, which is then converted to a more stable 4-carbon organic acid, malate or Asp, which diffuse to the bundle sheath cells (Gowik and Westhoff, 2011). Here the C<sub>4</sub> acid is decarboxylated, releasing CO<sub>2</sub> near the active site of Rubisco, which is located only in this cell type in C<sub>4</sub> plants. Given the higher efficiency of the C<sub>4</sub> photosynthetic mechanism under current atmospheric [CO<sub>2</sub>], efforts are underway to install C<sub>4</sub> photosynthesis in C<sub>3</sub> plants such as rice (the International C<sub>4</sub> rice consortium, <http://c4rice.irri.org/>) and other crops ([www.3to4.org](http://www.3to4.org)). While the number of genes necessary for the main enzymatic reactions and transporters involved in C<sub>4</sub> photosynthesis is relatively small, the introduction of C<sub>4</sub> photosynthesis into C<sub>3</sub> crops will also require major changes in leaf anatomy (von Caemmerer *et al.*, 2012). Initial progress toward the identification of the genes responsible for C<sub>4</sub> anatomy has been reported (Feldman *et al.*, 2014; Rizal *et al.*, 2015). On the other hand, terrestrial plants capable to carry out C<sub>4</sub> photosynthesis within a single cell were discovered about 10 years ago (Sharpe and Offermann, 2014). While these plants lack the typical Kranz features, they possess a subcellular separation that enables a concentrating of CO<sub>2</sub> at the active site of Rubisco. The genes involved in the development of this peculiar subcellular anatomy are unknown. Considering the scarcity of sequence information for single cell C<sub>4</sub> species, it is difficult to judge if single cell C<sub>4</sub> metabolism can be bio-engineered into C<sub>3</sub> crops.

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#### 198 *Introduction of CO<sub>2</sub>-concentrating mechanisms into chloroplasts*

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200 Another strategy to reduce oxygenation and thereby photorespiration is to  
201 introduce cyanobacterial CO<sub>2</sub>-concentrating mechanisms (CCM) into the  
202 chloroplasts of land plants (Zarzycki *et al.*, 2013). Cyanobacteria suppress the

oxygenating reaction of Rubisco by concentrating CO<sub>2</sub> inside a proteinaceous microcompartment called carboxysome. The  $\beta$ -carboxysome is constituted by an outer shell composed of several different proteins that enclose Rubisco and carbonic anhydrase, which releases CO<sub>2</sub> inside the carboxysome. The high [CO<sub>2</sub>] obtained near to the active site of cyanobacterial Rubisco suppresses oxygenation thereby increasing the catalytic efficiency of the carboxylation reaction of the enzyme. Furthermore, the use of CCM paves the way to potentially replace the native Rubisco with the cyanobacterial enzyme that has higher catalytic rate but also a lower affinity for CO<sub>2</sub> and specificity factor (meaning that is more prone to oxygenating RuBP) compared to the plant one (Zarzycki *et al.*, 2013). This would reduce the amount of Rubisco needed to sustain photosynthesis and permit the allocation of nitrogen for other purposes, thus increasing nitrogen use efficiency (Zhu *et al.*, 2004). The feasibility of introducing carboxysomes into higher plants was boosted by Lin *et al.*, (2014a) demonstration that the shell proteins of the  $\beta$ -carboxysome could be assembled in *Nicotiana benthamiana* chloroplasts producing organized, although empty, microcompartments. The same group was also able to introduce a functional cyanobacterial Rubisco in tobacco chloroplasts together with an internal carboxysomal protein (Lin *et al.*, 2014b). In this instance they replaced the native *Nicotiana tabacum* gene encoding for the large subunit of Rubisco and replaced it with the large and small subunits of the *Synechococcus elongatus* Rubisco, an enzyme with lower CO<sub>2</sub> affinity but higher catalytic rate compared to the endogenous one. The transformed lines were photosynthetically competent albeit at very high [CO<sub>2</sub>] and the formation of complexes between the cyanobacterial Rubisco and the carboxysomal protein was observed within the chloroplast stroma as occurs during cyanobacterial  $\beta$ -carboxysomes biogenesis, representing an important step toward the introduction of a CCM into C<sub>3</sub> plants. Simpler CCM mechanisms have been also considered for the transformation of C<sub>3</sub> plants. For example, a recent work described the introduction of a cyanobacterial bicarbonate transporter into tobacco chloroplasts (Pengelly *et al.*, 2014). The transformed plants expressed ample amount of the foreign transporter but displayed the same CO<sub>2</sub>-assimilation rates than the WT, implying that the transporter had little or no *in vivo* activity.

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236 *Rubisco engineering and screening for natural variation*



237

238 Despite its central role in plant metabolism, Rubisco is a relatively inefficient  
239 enzyme (Carmo-Silva *et al.*, 2014). In addition to its oxygenase activity, Rubisco  
240 also shows a relatively low  $k_{cat}$  value for CO<sub>2</sub> that obliges plants to produce very  
241 high amounts of the enzyme in order to sustain adequate photosynthesis,  
242 representing a large nitrogen investment (Zhu *et al.*, 2007). Understandably,  
243 considerable effort has been made to address these inefficiencies by trying to  
244 engineer a more efficient Rubisco. One first challenge for replacing the plant  
245 endogenous Rubisco with a more efficient one is that the large subunit of the  
246 enzyme is encoded by a single chloroplastic gene and the small one by several  
247 nuclear genes. Transformation of both the nuclear and chloroplast genomes of the  
248 same plant is thus required in order to substitute the endogenous enzyme with a  
249 more efficient one. Given that the active sites of Rubisco are on the chloroplast-  
250 encoded large subunit (Andersson, 2008), it may be possible that changing only  
251 the large subunit will improve enzyme efficiency, but this would require the  
252 transformation of the chloroplast genome, a technique that is currently available  
253 only for a small number of species. High-resolution crystallographic structural  
254 data are available for several plant Rubiscos and were used in site-directed  
255 mutagenesis approaches in order to try to improve Rubisco efficiency. However,  
256 this effort was hindered by the propensity of plant Rubisco to form insoluble  
257 aggregates when expressed in *E. coli*, probably caused by the lack of the complex  
258 network of chaperonins needed for the correct folding of the plant enzyme in the  
259 bacterial host (Saschenbrecker *et al.*, 2007; Liu *et al.*, 2010; Feiz *et al.*, 2012).  
260 For this reason, structure-function studies were carried out mainly with the  
261 enzymes from cyanobacteria and from the alga *Chlamydomonas reinhardtii*  
262 (Whitney *et al.*, 2011a; Parry *et al.*, 2013 and references therein). Another  
263 limitation to rational Rubisco engineering is our poor knowledge of the  
264 mechanism of Rubisco-catalysed oxygenation (Tcherkez, 2015). To overcome  
265 these technical difficulties, Whitney *et al.* (2011b) used transplastomic tobacco  
266 lines that expressed WT and mutated genes encoding the large Rubisco subunit  
267 from either C<sub>3</sub> or C<sub>4</sub> plants as well as from C<sub>3</sub>-C<sub>4</sub> intermediate species. Using this  
268 approach, the investigators were able to identify a single amino acid residue  
269 responsible for the different catalytic properties of the Rubiscos from C<sub>3</sub> and C<sub>4</sub>  
270 plants (low  $k_{cat}$  combined with low  $K_m$  for CO<sub>2</sub> and high  $k_{cat}$  combined with high

271  $K_m$  for CO<sub>2</sub>, respectively). Together, these results have opened the door to further  
272 possibilities for crop improvement. In fact, the co-engineering of a C<sub>4</sub>-type  
273 Rubisco with high  $k_{cat}$  for CO<sub>2</sub> together with the engineering of a CCM in the  
274 chloroplast to compensate for its low affinity for CO<sub>2</sub> may in theory be able to  
275 greatly enhance C<sub>3</sub> plant yield. Even without engineering CCM into chloroplasts,  
276 the raise in CO<sub>2</sub> levels that is expected by the end of the century will also  
277 probably allow for a less specific, and hence faster Rubisco. More complex  
278 approaches for the optimization of Rubisco via the manipulation of the activation  
279 state of the enzyme and its interaction with the various effectors that modulate its  
280 activity can also be envisaged (see the review of Carmo-Silva *et al.*, 2014).

281         The enormous natural variability that exists between terrestrial plants can  
282 be exploited in order to develop new strategies for reducing photorespiratory  
283 losses. Plants have developed several strategies, both anatomical and metabolic, to  
284 reduce photorespiration and compensate for its inhibitory effects (Sage, 2013).  
285 However, several of these mechanisms such as the regulation of leaf temperature,  
286 regulation of stomatal opening, establishment of CCM etc. are generally  
287 controlled by large sets of genes, some of which are unknown. On the other hand,  
288 Rubisco is encoded by a small set of known genes and the natural variability of  
289 this enzyme among different plant species has been taken into consideration in  
290 order to look for more efficient forms of the enzyme. The Rubisco specificity  
291 factor (i.e. the ratio of carboxylation to oxygenation at any given ratio of [CO<sub>2</sub>]  
292 and [O<sub>2</sub>]) displays some variation among the different C<sub>3</sub> species. For example,  
293 species growing in hot and dry environments seem to have Rubiscos with higher  
294 specificity factor (Galmés *et al.*, 2005), which may be taken into consideration as  
295 a criteria for selection of candidates to use in the substitution of the less efficient  
296 endogenous enzymes of different C<sub>3</sub> crops. While the potential of more efficient  
297 forms of Rubisco has yet to be exploited, several theoretical models suggest that  
298 changing the endogenous Rubisco with an enzyme with a more favourable  
299 specificity factor may improve crop yields (Zhu *et al.*, 2004; Parry *et al.*, 2011). It  
300 should be also taken into consideration that the Rubisco specificity factor may not  
301 necessarily reflect the effectiveness of the enzyme depending on the mechanism  
302 of the oxygenation reaction, which is still not completely known (Tcherkez,  
303 2015).

304           The natural variability of photorespiration is not only limited to the  
305 variation in the characteristics of Rubisco. Species-specific changes in the route  
306 are also possible, which implies that the pathway may be different from the basic  
307 “textbook” version. For example, it was demonstrated that the conversion of  
308 hydroxypyruvate to glycerate can also occur in the cytosol (Timm *et al.*, 2008).  
309 Arabidopsis may also show peculiar characteristics in the reassimilation of  
310 photorespiratory  $\text{NH}_3$ . In fact, mutants of plastidic GS<sub>2</sub>, the enzyme in charge of  
311 the reassimilation of photorespiratory ammonium, have been isolated in barley  
312 (Blackwell *et al.*, 1988) and in the model legume *Lotus japonicus* (Pérez-Delgado  
313 *et al.*, 2013) by screening an EMS population for the typical “photorespiratory”  
314 phenotype. However, no plastidic GS<sub>2</sub> mutants have been found in Arabidopsis.  
315 Given that the mutagenesis screen that was carried out with these plants was  
316 probably saturating (for example, 58 mutants were found affecting Fd-GOGAT,  
317 the other plastidic enzyme involved in  $\text{NH}_3$  reassimilation) and that Arabidopsis  
318 GS<sub>2</sub> is encoded, as in most plants, by a single gene (At5g35630), it is puzzling  
319 why GS<sub>2</sub> mutants were not been isolated either in the original screening or by  
320 means of transposon insertion. Another example of variation in photorespiratory  
321 metabolism related to ammonia reassimilation can be found in conifers, where the  
322 plastidic isoform of GS is not present but, unlike other higher plants, a cytosolic  
323 GS isoform is expressed in photosynthetic cells, and photorespiratory ammonia is  
324 probably reassimilated through a cytosolic GS/GOGAT cycle (Ávila *et al.*, 2001).

325

#### 326 *Photorespiratory bypasses*

327

328       Instead of trying to reduce the photorespiratory rates, a different approach is to  
329 install alternative and less energetically expensive routes for the recycling of  
330 2PG. Three bypasses to the reactions of the photorespiratory pathway were  
331 successfully engineered in Arabidopsis. In the first approach, glycolate was  
332 converted to glycerate directly in the chloroplast by introducing the *Escherichia*  
333 *coli* glycolate catabolic pathway, thus avoiding or at least competing with the  
334 peroxisomal and mitochondrial reactions of photorespiration (Kebeish *et al.*,  
335 2007). The second approach was to introduce a complete glycolate catabolic  
336 cycle that oxidized 2PG to  $\text{CO}_2$  in the chloroplast (Maier *et al.*, 2012). Both  
337 bypasses should avoid ammonia release in the mitochondria, which is quite

expensive to reassimilate in terms of the ATP and reducing equivalents required. However, while the “Kebeish” bypass resulted in an improved energy balance, the “Maier” bypass was costlier compared to the standard photorespiratory cycle (Peterhänzel *et al.*, 2013b). Despite this, both bypasses were reported to enhance biomass production by up to 30%. In the case of the “Maier” bypass it is speculated that this benefit may be due to the release of CO<sub>2</sub> from 2PG oxidation directly in the chloroplast, this might increase the chloroplastic CO<sub>2</sub> concentration and reduce the probability of further oxygenating reactions. Interestingly, both bypasses resulted in increased biomass production only under short-day conditions but not in long days. A third bypass to photorespiration has been engineered by introducing the *E. coli* enzymes glyoxylate carboligase and hydroxypyruvate isomerase into tobacco for the conversion of glyoxylate into hydroxypyruvate directly in the peroxisome, thus once again avoiding ammonia release in the mitochondria (Carvalho *et al.*, 2011). While this alternative pathway may potentially reduce the cost of 2PG recycling (Peterhänzel *et al.*, 2013b), hydroxypyruvate isomerase protein was not detectable in these tobacco lines, so its impact on plant yield remains to be proven. In recent reports, the potential of photorespiratory bypasses for the improvement of plants of agronomical importance has been demonstrated. It was shown that introduction of the “Kebeish” bypass in the oilseed crop *Camelina sativa* greatly increased seed yield, which may be used for the production of biofuels (Dalal *et al.*, 2015). Also, in another study, potato (*Solanum tuberosum*) plants were transformed with the three genes that encode for *E. coli* glycolate dehydrogenase subunits and the corresponding polyprotein was successfully expressed in the chloroplast, where it was able to catalyze the conversion of glycolate to glyoxylate (Nölke *et al.*, 2014). The enhancement in assimilation rate led to an increase in shoot biomass and subsequently to a greater tuber yield in the transgenic lines. This suggested that part of the glyoxylate produced in the chloroplast by the bacterial enzyme may be completely oxidized *in situ* to CO<sub>2</sub> that would be released near the Rubisco active site and would thereby reduce the rate of Rubisco oxygenation. Recent evidences support the idea that glyoxylate can be decarboxylated in the chloroplast by the action of the endogenous pyruvate dehydrogenase (Blume *et al.*, 2013). However, in order to try to establish a highly efficient partial or complete ‘Kebeish’ bypass, it should be taken into consideration that plastids

372 contain a highly active NADPH-dependent glyoxylate dehydrogenase, which is  
373 able to reduce this molecule back to glycolate (Allan *et al.*, 2009) and should be  
374 probably silenced in order to avoid a futile cycle in the chloroplast.

375 Completely new bypasses can be also designed by taking advantage of the  
376 enormous amount of different enzyme activities that can be found in bacteria,  
377 algae and Archeae (see Ort *et al.*, 2015 for some examples). More ambitious  
378 approaches would be to design bypasses that involve intermediates that are not  
379 present in the plant or to genetically engineer a single enzyme able to degrade  
380 2PG to CO<sub>2</sub> directly in the chloroplast. In a recent report, a synthetic pathway  
381 that worked both as a photorespiratory bypass and as an additional CO<sub>2</sub>-fixing  
382 pathway, the hydroxypropionate bi-cycle was successfully engineered in a  
383 cyanobacterium (Shih *et al.*, 2014). Simulated energy balance analyses can be  
384 performed in order to predict the potential benefits of a bypass to photorespiration  
385 (Xin *et al.*, 2015).

386 When designing synthetic routes for the recycling of 2PG, it has to be  
387 taken into consideration that alternative routes to the core photorespiratory  
388 pathway are already present in nature, although their physiological meaning and  
389 the flux that may pass through them is not known. For example, glyoxylate can  
390 be oxidatively decarboxylated to formate and CO<sub>2</sub> probably by a non-enzymatic  
391 reaction that takes place in the peroxisomes of higher plants in the presence of  
392 H<sub>2</sub>O<sub>2</sub> (Igamberdiev *et al.*, 1999). Cyanobacteria on the other hand are able to  
393 enzymatically decarboxylate glyoxylate via oxalate by using an alternative  
394 pathway for the recycling of 2PG (Eisenhut *et al.*, 2008). In barley mutants with  
395 reduced glycine decarboxylase (GDC) activity, this formate may be used to  
396 support the synthesis of serine through a GDC-independent pathway that does not  
397 release NH<sub>3</sub>, thus greatly reducing the energy cost of the photorespiratory cycle  
398 (Wingler *et al.* 1999a). As aforementioned, glyoxylate can be decarboxylated in  
399 the chloroplast by the action of the endogenous pyruvate dehydrogenase (Blume  
400 *et al.*, 2013), and a cytosolic hydroxypyruvate reductase provides an alternative  
401 route to the peroxisomal conversion of hydroxypyruvate to glycerate (Timm *et al.*  
402 *et al.*, 2008). Several other possibilities for peroxide-mediated decarboxylations  
403 have also been proposed (Grodzinski and Butt 1977; Cousins *et al.* 2008; Keech  
404 *et al.* 2012), but the extent to which these reactions would happen under natural  
405 conditions still remains unclear. Therefore, a current challenge resides in finding

406 better tools to challenge these alternative pathways and assess their natural  
407 occurrence under both normal and stress conditions.

408

409 *Optimization of the levels of photorespiratory enzymes*

410

411 Analysis of dynamic metabolic models of photosynthetic carbon metabolism  
412 suggested that there may be an underinvestment of resources in the biosynthesis  
413 of Rubisco and of the enzymes of the Calvin-Benson cycle and concomitantly an  
414 overinvestment in photorespiratory enzymes. This scenario may be responsible of  
415 a less than optimal photosynthetic efficiency leading to reduced crop yields (Zhu  
416 *et al.*, 2007). Interestingly, this appears rather contradictory to recent studies in  
417 which the amount of photorespiratory enzymes has been modulated. For instance,  
418 different studies carried out in crops species indicate that antisense reduction of  
419 individual photorespiratory enzymes is associated with lower productivity. Potato  
420 plants with reduced levels of the GDC-P protein (Heineke *et al.*, 2001) or of  
421 serine hydroxymethyltransferase (Schjoerring *et al.*, 2006) as well as rice plants  
422 with lower levels of glycolate oxidase (Xu *et al.*, 2009) showed reduced  
423 photosynthetic and growth rates. By contrast, a few studies have reported an  
424 improved performance of plants with increased levels of photorespiratory  
425 enzymes. Overexpression of GDC-H protein or the GDC-L protein in  
426 Arabidopsis resulted in enhanced net-photosynthesis and plant growth (Timm *et al.*,  
427 2012a; Timm *et al.*, 2015). Increased yields were not observed under elevated  
428 CO<sub>2</sub> atmosphere, indicating that they were due to a facilitated carbon flow  
429 through GDC and the photorespiratory pathway as a whole. It is assumed that  
430 increased photorespiratory capacity may reduce negative feedback exerted by  
431 photorespiratory metabolites on the Calvin-Benson cycle thus enhancing CO<sub>2</sub>  
432 assimilation. Recent data suggest that 2PG levels could be of key importance in  
433 this coordinated control of photosynthesis and photorespiration (Timm *et al.*,  
434 2012b; Haimovich-Dayana *et al.*, 2015). Overexpression of serine  
435 hydroxymethyltransferase, the enzyme that acts in conjunction with glycine  
436 decarboxylase to produce serine in the mitochondrion, was also able to improve  
437 photosynthetic efficiency and plant productivity in rice (Wu *et al.*, 2015). Taken  
438 together, these results clearly indicate that the mitochondrial conversion of  
439 glycine to serine is a bottleneck of the photorespiratory pathway or is somehow

otherwise involved in the regulation of photosynthetic activity. The recent discovery that serine may act as a metabolic signal for the transcriptional regulation of photorespiration (Timm *et al.*, 2013) further supports this idea. In addition to the reactions involved in the glycine to serine conversion, the reassimilation of photorespiratory  $\text{NH}_4^+$  is probably another bottleneck of the photorespiratory pathway. Photorespiratory  $\text{NH}_4^+$  is reassimilated by the action of the plastidic isoform of glutamine synthetase ( $\text{GS}_2$ ), and it has been suggested that this reaction must be the rate-limiting step of the pathway (Wallsgrave *et al.*, 1987, Häusler *et al.*, 1994; Kozaki and Takeba, 1996; Hoshida *et al.*, 2000). Plants that overexpress  $\text{GS}_2$  showed enhanced growth rate under active photorespiratory conditions (Migge *et al.*, 2000; Zhu *et al.*, 2014). Unfortunately, the growth of these  $\text{GS}_2$  overexpressors was compared to WT plants under normal air conditions but not under  $\text{CO}_2$ -enriched atmosphere, so it cannot be ruled out if the increased yield is due to improved nitrogen assimilation rather than to an increased capacity for photorespiration (Migge *et al.*, 2000; Zhu *et al.*, 2014). However, the fact that mutants lacking  $\text{GS}_2$  show a similar growth rate compared to wild-type plants under photorespiratory-suppressed conditions (Wallsgrave *et al.*, 1987; Betti *et al.*, 2014) indicates that  $\text{GS}_2$  is not probably playing an important role in primary nitrogen assimilation. Moreover, overexpression of  $\text{GS}_2$  confers resistance under stress conditions like salinity or high light (Kozaki and Takeba, 1996; Hoshida *et al.*, 2000). Taking into consideration the promising results obtained with these overexpressors, it would be also worth to exploit natural variability and look for cultivars that already have higher or lower levels of photorespiratory enzymes.

Another important and often neglected parameter lies in the transcriptional and post-translational modifications of photorespiratory genes and enzymes. Different reports suggest that at the transcriptional level photorespiratory genes are regulated in a similar way to the photosynthetic ones (Foyer *et al.*, 2009; Pérez-Delgado *et al.*, 2013). On the other hand, metabolic data analysis of WT and photorespiratory mutants under different  $\text{CO}_2$  and  $\text{O}_2$  conditions suggest a fine tuning of photorespiratory metabolism (Timm *et al.*, 2012b). Regarding post-translational modifications, it was recently shown that seven enzymes of the photorespiratory cycle could be phosphorylated (Hodges *et al.*, 2013). Furthermore, looking to redox proteome data, it appeared that almost all

474 photorespiratory enzymes could undergo oxidative modifications for some of  
475 their cysteine residues, and were therefore identified as potential targets for redox  
476 regulations (Keech *et al.*, submitted for publication). Undoubtedly, the next step  
477 will be to determine primarily the extent to and the conditions for which the  
478 proteins or cysteines are modified, the type of modifications that occur, and  
479 secondly whether these modifications positively or negatively regulate enzyme  
480 activities, and how they are controlled at the cellular level. Altogether, this  
481 clearly indicates that a rational bio-engineering of plants with modified levels of  
482 photorespiratory enzymes would also benefit from an increased knowledge of the  
483 biochemical regulations inherent to this cycle.

484

#### 485 *Perspectives for crop improvement*

486

487 As summarized in the above sections and in Table 1, several approaches have  
488 been used in order to manipulate photorespiration in attempt to increase plant  
489 yield. However, most of these efforts have been carried out using model plants  
490 (with some notable exceptions like the consortia working on the transformation of  
491 rice into a C<sub>4</sub> plant, see <http://c4rice.irri.org/>). In the light of the results obtained  
492 by recent field trials (Aliyev, 2012), it would appear unlikely that crops with  
493 improved photorespiratory performance can be obtained by screening for natural  
494 genetic variation, but they should be rather generated by means of genetic  
495 engineering. Unfortunately, transformation of our major crops is still a difficult  
496 and time-consuming process, even if is getting easier and more successful every  
497 year. Moreover, some promising approaches such as the engineering of the large  
498 subunit of Rubisco require the transformation of chloroplast DNA, a technique  
499 that is available only for a few crop species: notably tobacco, potato, tomato and  
500 perhaps soybean, but as yet not cereal species (Scharff and Bock, 2014). As a first  
501 step, organisms for which transformation is more tractable such as algae and  
502 cyanobacteria can be used in order to obtain clues on the metabolic and  
503 physiological consequences of a targeted genetic manipulation. A second step  
504 may be the use of tobacco; a plant that is especially easy to transform both in the  
505 nuclear and plastid genomes and forms canopies in the field that are similar to  
506 those of food crops (Long *et al.*, 2015). Even after careful experimental design  
507 and test in intermediate plant models, several challenges would need to be



508 overcome before new genes and pathways can be introduced into crops. As  
509 mentioned before, nuclear and especially plastid transformation techniques are  
510 still inefficient or unavailable for most staple crops. In addition to that, promoters  
511 and vectors that can permit high expression of transgenes and a correct subcellular  
512 localization of the protein product should be available, together with strategies to  
513 avoid gene silencing and random insertion in the genome (see Ort *et al.*, 2015 for  
514 a more detailed discussion on this topic). It should also be taken into consideration  
515 that crops with engineered photorespiratory pathways will be considered as  
516 genetically modified plants (GMP), and the potential use of such GMPs will  
517 remain limited under the current legislation, which furthermore can vary greatly  
518 between countries. For example in the European Union the authorization  
519 procedure for placing a GMP on the market is a long, complex and expensive  
520 procedure regulated by directives that were approved more than 10 years ago  
521 (more details in Hartung and Schiemann, 2014). Furthermore, due to social and  
522 political rejection of GMPs, even those transgenic plants that have been approved  
523 are not cultivated in most EU countries. On the other hand, several millions of  
524 hectares of GMPs are growing in countries with less restrictive regulations such as  
525 the United States, Canada, Brazil, India and China. That said, several new  
526 molecular techniques, like TALENS (transcription activator-like effector  
527 nuclease(s)) or the CRISPR/Cas9 system, have been developed in the recent  
528 years. The use of these genome editing techniques can lead to the production of  
529 plants which cannot be classified as GMPs under current legislations. The  
530 European Commission is currently evaluating these techniques together with  
531 cisgenesis and intragenesis, RNA-dependent DNA methylation, grafting  
532 (production of chimeric plant with a wild-type scion inserted on a genetically  
533 modified rootstock), reverse breeding and agro-infiltration in order to determine  
534 the extent to which they should lead to genetically modified organisms (Lusser *et*  
535 *al.*, 2012). Promising steps towards the regulation of these techniques are being  
536 given, for example mutant plants obtained with the CRISPR/Cas9 system have not  
537 been considered as GMPs in a recent decision of the Swedish Board of  
538 Agriculture ([http://www.upsc.se/about-upsc/news/4815-green-light-in-the-tunnel-](http://www.upsc.se/about-upsc/news/4815-green-light-in-the-tunnel-swedish-board-of-agriculture-a-crispr-cas9-mutant-but-not-a-gmo.html)  
539 [swedish-board-of-agriculture-a-crispr-cas9-mutant-but-not-a-gmo.html](http://www.upsc.se/about-upsc/news/4815-green-light-in-the-tunnel-swedish-board-of-agriculture-a-crispr-cas9-mutant-but-not-a-gmo.html)).

540

541 *Should we really look for plants with lower rates of photorespiration?*

542

543 Regardless of the difficulties that we may face to obtain plants with modified  
544 photorespiratory rates, some changes in photorespiration in the field will happen  
545 anyway because of the rise in atmospheric [CO<sub>2</sub>], which is predicted to double by  
546 2100 (Intergovernmental Panel on Climate Change, 2014). On one hand, this  
547 increase in [CO<sub>2</sub>] will reduce photorespiration by increasing CO<sub>2</sub> fixation by  
548 Rubisco. On the other hand, photorespiration should be stimulated by the  
549 predicted increase of the average atmosphere temperature, and subsequently of  
550 leaf canopy. Moreover, the expected increased stomatal closure caused by  
551 elevated CO<sub>2</sub> will contribute to further increase in leaf temperature. Thus,  
552 photorespiratory losses are still expected to be high even in a high CO<sub>2</sub> world.  
553 Photorespiration has been traditionally considered as a wasteful and unavoidable  
554 process that needs to be minimized in order to improve plant yield. However,  
555 different lines of evidence suggest that reducing photorespiration may not  
556 necessarily always have beneficial effects.

557 1) Plant productivity may be improved by engineering more efficient ways to  
558 recycle 2PG but also by an increased capacity for photorespiratory flux. The  
559 introduction of bypasses to photorespiration can lead to up to 30% of increase in  
560 plant biomass (Kebeish *et al.*, 2007; Maier *et al.*, 2012; Nölke *et al.*, 2014).  
561 However, these beneficial effects were observed only under short day conditions  
562 and/or controlled temperature and humidity, which may not always reflect the  
563 conditions that crops will face in the field. Further testing of these GMPs under  
564 different conditions would be needed in order to determine if photorespiratory  
565 bypasses may be beneficial also under field conditions. By contrast, several  
566 studies indicated that a higher capacity for photorespiratory flux is paralleled by  
567 increased plant yield (see the section “Optimization of the levels of  
568 photorespiratory enzymes”). A higher photorespiratory capacity would reduce the  
569 levels of photorespiratory metabolites that may inhibit the Calvin-Benson cycle as  
570 well as increase the rate at which photorespiratory carbon is returned to the  
571 chloroplast in form of 3-PGA, thus facilitating CO<sub>2</sub> assimilation. Therefore, CO<sub>2</sub>  
572 assimilation may be improved either by bypassing photorespiration or by the  
573 overexpression of bottleneck enzymes of the cycle. The best engineering strategy  
574 to use will depend on the crop considered and the environmental conditions at the  
575 field level.

2) Energetically wasteful and useful are not necessarily antithetic to one another. As mentioned before, under stress conditions such as drought, salinity, cold, high light, heat or a combination of them, an excess of NADPH may be produced that could lead to an increase of reactive oxygen species (ROS). Photorespiration can act as a sink for this excess of reducing power, and this welcome effect can be even more important considering that different stress conditions can increase photorespiratory rates. Drought and salinity for example trigger a decrease in stomatal conductance, thus decreasing the  $\text{CO}_2:\text{O}_2$  ratio and increasing photorespiration (Kangasjärvi *et al.*, 2012). Heat also leads to increased photorespiration of decreased Rubisco specificity and secondarily due to the changes in the relative solubility of  $\text{CO}_2$  and  $\text{O}_2$ . It is not surprising then that attention has been paid to the role of photorespiration in the response to stress (Wingler *et al.*, 2000; Voss *et al.*, 2013). Barley mutants with reduced levels of different photorespiratory enzymes as well as Arabidopsis mutants of the peroxisomal hydroxypyruvate reductase (HPR1) enzyme were more sensitive to drought (Wingler *et al.*, 1999b; Li and Hu, 2015). On the other hand, rice plants with increased photorespiratory capacity showed enhanced tolerance to salt stress (Hoshida *et al.*, 2000). A protective role of photorespiration in the dissipation of excess energy has been already hypothesized long time ago (Heber and Krause, 1980) and a demonstration to this hypothesis was provided later by Kozaki and Takeba (1996), who showed that photorespiration protects against photoinhibition caused by high light. A more recent work demonstrated that when the photorespiratory cycle is impaired, the excess of reducing power and the consequent over-production of ROS prevent the repair of photosystem II, thus leading to accelerated photoinhibition (Takahashi *et al.*, 2007). A role for photorespiration in the response to other kinds of stress such as chilling or exposure to heavy metals has also been proposed (Voss *et al.*, 2013 and references therein). Interestingly, several photorespiratory genes are co-expressed with genes involved in the resistance to Al, that although not technically a heavy metal is also a stressor that constrains plant productivity (Nunes-Nesi *et al.*, 2014a). Since abiotic stress is one of the factors that most frequently limits crop productivity worldwide (Mittler, 2006), the performance of plants with reduced rates of photorespiration should be tested carefully under different stress conditions. This should be carried out also for plants expressing bypasses to photorespiration, since

610 the sink effect for excess reducing power exerted by photorespiration under stress  
611 conditions may be lost in such organisms. Moreover, since most of the high  
612 quality soils available are already farmed, the rising demand for food would  
613 probably lead to farm crops in marginal lands with poorer soil and adverse  
614 climatic conditions. In such a scenario, the use of crops with high resistance to  
615 abiotic stress, and not only high yield under optimal conditions, would seem to be  
616 desirable.

617 Interestingly, photorespiration has also been shown to play a significant  
618 role in biotic stress responses, where the  $\text{H}_2\text{O}_2$  produced by the reaction of  
619 glycolate oxidase in the peroxisome plays a central role in the defence from  
620 pathogen attack (Taler *et al.*, 2004; Rojas *et al.*, 2012) and is part of the signalling  
621 route that leads to programmed cell death (Mateo *et al.*, 2004). Plants with  
622 reduced rates of photorespiration or engineered with alternative routes that bypass  
623 the peroxisomal part of the pathway may show increased sensitivity to pathogen  
624 attacks and should also be tested carefully. In a recent report it was also showed  
625 that some photorespiratory enzymes are highly expressed in plant roots (Nunes-  
626 Nesi *et al.*, 2014b), so it is possible that changes in the levels of photorespiratory  
627 enzymes may also affect the physiology of heterotrophic tissues.

628 3) Rates of photorespiration correlate with nitrate assimilation in hydroponically  
629 grown *Arabidopsis* and wheat (Rachmilevitch *et al.*, 2004; Bloom *et al.*, 2010).  
630 This relationship has even been proposed to explain the lower-than-expected  
631 growth increases in plants grown under elevated  $\text{CO}_2$  and explain why many  $\text{C}_3$   
632 crops and trees grow more slowly when fed with nitrate as a sole nitrogen source  
633 (Bloom *et al.*, 2011). Recent evidence suggests that these hydroponic-based  
634 observations may occur at larger scales when it was shown that wheat grown  
635 under free-air  $\text{CO}_2$  enrichment had higher nitrate pools and a greater  $^{15}\text{N}$   
636 enrichment of both total nitrogen and nitrate, observations consistent with a  
637 decrease in nitrate assimilation (Bloom *et al.*, 2014). The exact mechanism that  
638 underpins this co-dependency is still unknown but it may be related to the  
639 photosynthesis-dependent export of malate from the chloroplast (the ‘malate  
640 valve’), which increases the levels of cytosolic NADH thus providing reducing  
641 equivalents for nitrate reduction (Bloom *et al.*, 2010). Additionally, increased  
642 rates of photorespiration further result in excess NAD(P)H since photorespiration  
643 consumes more ATP relative to NAD(P)H than  $\text{CO}_2$  fixation (Kramer and Evans,

2011; Walker et al., 2014). This results in excess NAD(P)H that must be consumed to balance the energy demands of central metabolism with energy production from the light reactions. C<sub>4</sub> plants on the other hand assimilate NO<sub>3</sub><sup>-</sup> independently of atmospheric CO<sub>2</sub> concentration since the cytoplasmic NADH for nitrate reduction can be produced by the same C<sub>4</sub> pathway instead of by photorespiration (Bloom, 2015).

Nitrate is the most abundant form of N in agricultural soils and is the major N source for most higher plants. This is despite the higher amount of energy that is needed for the assimilation of NO<sub>3</sub><sup>-</sup> into organic compounds compared to other N sources such as NH<sub>4</sub><sup>+</sup> or organic forms of nitrogen. Taking this into consideration, it is possible that a reduction of the photorespiratory rates in crops that use mainly NO<sub>3</sub><sup>-</sup> may lead to nitrogen deprivation. Reliance on NH<sub>4</sub><sup>+</sup> fertilizers may not always be possible in order to circumvent this since many plants show symptoms of toxicity when grown on NH<sub>4</sub><sup>+</sup> as the sole N source (Britto and Kronzucker, 2002).

In conclusion, different lines of evidence have shown that engineering of photorespiration may greatly improve plant CO<sub>2</sub>-assimilation and growth. Several recent advances have been made in reducing photorespiratory losses in model organisms as well as in some plants of agricultural relevance. A great challenge will be the transfer of these advances to our major food crops, which are generally more recalcitrant to genetic manipulation. Nonetheless, a rational bio-engineering of plants with altered photorespiration should also take into consideration that this pathway is tightly connected with several other aspects of plant metabolism and a reduction of photorespiration may not always be beneficial, especially for plants growing under adverse environmental conditions. Finally, taking into consideration that NO<sub>3</sub><sup>-</sup> assimilation depends on photorespiration, the manipulation of the photorespiratory pathway may also affect the rates of N assimilation and may favour the use of one N source over another.

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**Table 1.** *Summary of the approaches that can be used to improve crop yield through manipulation of photorespiration.*

| Strategy   | Advantages (A) / Disadvantages (D)  |
|--|---|
| Screening for plants with reduced rates of photorespiration                      | A: -No need for genetic manipulation.<br>D: -Highly improbable to find plants with high levels of PS and low of photorespiration in field trials.   |
| Enhancing the amount of photorespiratory CO <sub>2</sub> scavenging              | A: -Does not imply changes in cellular metabolism.<br>D: -Genetic determinants of organelle partitioning and connection are not completely understood.  |
| Introduce C <sub>4</sub> photosynthesis into C <sub>3</sub> plants               | A: -Great theoretical potential for increase in crop yield.<br>D: -Major changes in leaf anatomy are required. -The genes responsible for C <sub>4</sub> anatomy not completely identified.   |
| Introduction of CCM into chloroplasts  | A: -Should greatly reduce the rates of photorespiration.<br>-Should allow replacing endogenous Rubisco with enzymes with higher catalytic rates and lower CO <sub>2</sub> affinity.<br>D: -Requires transformation of the chloroplast genome.<br>-Complex CCM requires the transformation of multiple genes and the correct assembly of multiprotein complexes.   |
| Rubisco engineering and screening for naturally occurring more efficient Rubisco | A: -Rubisco has several catalytic inefficiencies. This implies several opportunities for engineering. -Naturally occurring more efficient Rubiscos have been found in some species.<br>D: -Structure-function studies with Rubisco are hampered by different technical difficulties. -The exact mechanism of the oxygenating reaction is still not completely understood.   |
| Photorespiratory bypasses  | A: -Successfully engineered in both model and crop species. -Can increase yield up to 30%. -Possibility of complete oxidation of 2PG in the chloroplast, thus raising the [CO <sub>2</sub> ] near Rubisco active site.<br>D: -Need transfer of multiple genes. -Increased yield is seen only under short day in some bypasses. -The lower energy cost of some bypasses may prevent the protective role of PR under stress conditions. |
| Optimization of the levels of photorespiratory enzymes                           | A: -Relatively easy genetic manipulation.<br>D: -Transcriptional and post-translational regulation of photorespiratory genes and enzymes is still poorly characterised.   |